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Daytime formation of nitrous acid at a coastal remote site in Cyprus indicating a common ground source of atmospheric HONO and NO

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Abstract. Characterization of daytime sources of nitrous acid (HONO) is crucial to understand atmospheric oxidation and radical cycling in the planetary boundary layer. HONO and numerous other atmospheric trace constituents were measured on the Mediterranean island of Cyprus during the CYPHEX (CYprus PHotochemical EXperiment) campaign in summer 2014. Average volume mixing ratios of HONO were 35 pptv (± 25 pptv) with a HONO / NO_x ratio of 0.33, which was considerably higher than reported for most other rural and urban regions. Diel profiles of HONO showed peak values in the late morning (60 ± 28 pptv around 09:00 local time) and persistently high mixing ratios during daytime (45 ± 18 pptv), indicating that the photolytic loss of HONO is compensated by a strong daytime source. Budget analyses revealed unidentified sources producing up to 3.4×10^6 molecules cm⁻³ s⁻¹ of HONO and up to 2.0×10^7 molecules cm⁻³ s⁻¹ NO. Under humid conditions (relative humidity > 70 %), the source strengths of HONO and NO exhibited a close linear correlation ($R^2 =$ 0.72), suggesting a common source that may be attributable to emissions from microbial communities on soil surfaces.

1 Introduction

Nitrous acid (HONO) is an important component of the nitrogen cycle, being widespread in the environment. Either in its protonated form (HONO or HNO₂) or as nitrite ions (NO₂⁻) it can be found not only in the gas phase, on aerosol particles, in clouds and in dew droplets but also in soil, seawater and sediments (Foster et al., 1990; Rubio et al., 2002; Acker et al., 2005, 2008; Bianchi et al., 1997). It plays a key role in the oxidizing capacity of the atmosphere, as it is an important precursor of the OH radical, which initiates most atmospheric oxidations. OH radicals react with pollutants in the atmosphere to form mostly less toxic compounds (e.g. $CO + OH \rightarrow CO_2 + H_2O$; Levy, 1971). Volatile organic compounds (VOCs) react with OH, contributing to formation of secondary aerosols (SOAs), which can serve as cloud condensation nuclei (CCN; Arey et al., 1990; Duplissy et al., 2008). Furthermore OH oxidizes SO₂ to H₂SO₄, which condense subsequently to form aerosol particles (Zhou et al., 2013). In this way HONO has an indirect effect on the radiative budget and climate. In the first 2-3 h following sunrise, when OH production from other sources (photolysis of O₃ and formaldehyde) is relatively low, photolysis of HONO can be the major source of OH radicals as HONO concentrations may be high after accumulation during nighttime (Lammel and Cape, 1996; Czader et al., 2012; Mao et al., 2010). On average up to 30% of the daily OH budget in the boundary layer is provided by HONO photolysis (Alicke et al., 2002; Kleffmann et al., 2005; Ren et al., 2006), but it has been reported as high as 56% (Ren et al., 2003), with ambient HONO mixing ratios ranging from several parts per trillion by volume (pptv) in rural areas up to a few parts per billion by volume (ppbv) in highly polluted regions (Acker et al., 2006a, b; Costabile et al., 2010; Li et al., 2012; Michoud et al., 2014; Spataro et al., 2013; Su et al., 2008a; Zhou et al., 2002a).

In early studies, atmospheric HONO was assumed to be in a photostationary state (PSS) during daytime controlled by the gas-phase reaction of NO and OH (Reaction R1) and two loss reactions, which are the photolysis (Reaction R2) and the reaction with OH (Reaction R3).

$$OH + NO \rightarrow HONO$$
 (R1)

HONO $\xrightarrow{hv(300-405\,\text{nm})}$ OH + NO (R2)

$$HONO + OH \rightarrow NO_2 + H_2O \tag{R3}$$

However, field measurements in remote and rural locations as well as urban and polluted regions found severaltimes-higher daytime HONO concentrations than model predictions, suggesting a large unknown source (Kleffmann et al., 2003, 2005; Su et al., 2008a, 2011; Sörgel et al., 2011a; Michoud et al., 2014; Czader et al., 2012; Wong et al., 2013; Tang et al., 2015; Oswald et al., 2015) even after considering direct emission of HONO from combustion sources (Kessler and Platt, 1984; Kurtenbach et al., 2001). Heterogeneous reactions on aerosols have been proposed as an explanation for the missing source. The hydrolysis (Reaction R4; Finlayson-Pitts et al., 2003) and redox reactions of NO₂ have been intensively investigated on different kinds of surfaces such as fresh soot, aged particles or organic-coated particles (Ammann et al., 1998; Arens et al., 2001; Aubin and Abbatt, 2007; Bröske et al., 2003; Han et al., 2013; Kalberer et al., 1999; Kleffmann et al., 1999; Kleffmann and Wiesen, 2005; Lelievre et al., 2004). Minerals like SiO₂, CaCO₃, CaO, Al₂O₃ and Fe₂O₃ showed a catalytic effect on the hydrolysis of NO₂ (Kinugawa et al., 2011; Liu et al., 2015; Wang et al., 2003; Yabushita et al., 2009). Different kinds of surfaces (humic acid and other organic compounds, titanium dioxide, soot) can be photochemically activated, which leads to enhanced NO₂ uptake and HONO production (Reaction R5, George et al., 2005; Langridge et al., 2009; Monge et al., 2010; Ndour et al., 2008; Ramazan et al., 2004; Stemmler et al., 2007; Kebede et al., 2013). The photolysis of particulate nitric acid (HNO₃), nitrate (NO₃⁻) and nitro-phenols (R-NO₂) leads to HONO formation as well (Baergen and Donaldson, 2013; Bejan et al., 2006; Ramazan et al., 2004; Scharko et al., 2014; Zhou et al., 2003, 2011). But these reactions cannot account for the HONO levels observed during daytime (Elshorbany et al., 2012).

$$2NO_2 + H_2O \rightarrow HONO + HNO_3 \tag{R4}$$

surface
$$\xrightarrow{hv} e^- \xrightarrow{NO_2} NO_2^- \xrightarrow{H_2O} HONO + OH^-$$
 (R5)

On the other hand, soil nitrite, either biogenic or nonbiogenic, has been suggested as an effective source of HONO (Su et al., 2011; Oswald et al., 2013; Mamtimin et al., 2016). Depending on soil properties such as pH and water content and according to Henry's law, HONO can be released (Donaldson et al., 2014b; Su et al., 2011). This is consistent with field flux measurements showing HONO emission from the ground rather than deposition as is the case for HNO₃ (Harrison and Kitto, 1994; Kleffmann et al., 2003; Ren et al., 2011; Stutz et al., 2002; VandenBoer et al., 2013, 2014; Villena et al., 2011; Zhou et al., 2011). In a recent study, Weber et al. (2015) measured large HONO and NO emissions from dryland soils with microbial surface communities (so-called biological soil crusts). Many studies have shown decreasing HONO mixing ratios with altitude in the lowest few hundred meters of the troposphere, due to respective short atmospheric lifetime compared to vertical transport time (Wong et al., 2012, 2013; Vogel et al., 2003; VandenBoer et al., 2013, 2014; Zhang et al., 2009; Young et al., 2012). According to the modeling results of Wong et al. (2013), we estimate that the ground HONO source could be important for up to 200-300 m a.g.l. This indicates that HONO is more relevant for the OH budget close to the surface than in high-altitude air masses.

Several field studies also show a correlation of the unknown HONO source with solar radiation or the photolysis frequency of NO₂ J_{NO_2} (Su et al., 2008a; Soergel et al., 2011a; Wong et al., 2012; Costabile et al., 2010; Michoud et al., 2014; Oswald et al., 2015; Lee et al., 2016). This correlation can be explained either by the aforementioned photosensitized reactions or by temperature-dependent soilatmosphere exchange (Su et al., 2011). According to Su et al. (2011), the release of HONO from soil surfaces is controlled by both the soil (biogenic and chemical) production of nitrite and the gas-liquid-phase equilibrium. The solubility is strongly temperature-dependent, resulting in higher HONO emissions during noontime and high-radiation $J_{\rm NO_2}$ periods, and lower HONO emissions or even HONO deposition during the nighttime as further confirmed by VandenBoer et al. (2015). This temperature dependence exists not only for equilibrium over soil solution but also for adsorptiondesorption equilibrium over dry and humid soil surfaces (Li et al., 2016).

In this study we measured HONO and a suite of other atmospherically relevant trace gases in a coastal area on the Mediterranean island of Cyprus in summer 2014. Due to low local anthropogenic impact and low NO_x levels in aged air masses, but high solar radiation, this is an ideal site at which to investigate possible HONO sources and to gain a better understanding of HONO chemistry.

2 Instrumentation

HONO was measured with a commercial long-path absorption photometry instrument (effective light path 1.5 m, LOPAP, Quma, Wuppertal, Germany). LOPAP has a collecting efficiency of > 99 % for HONO and a detection limit of 4 pptv at a time resolution of 30 s. To avoid potential interferences induced by long inlet lines and heterogeneous formation or loss of HONO on the inlet walls (Kleffmann et al., 1998; Zhou et al., 2002b; Su et al., 2008b), HONO was collected by a sampling unit installed directly in the outdoor atmosphere, i.e., placed on a mast at a height of 5.8 m above ground installed at the edge of a laboratory container. Furthermore, the LOPAP has two stripping coils placed in series to reduce known interfering signals (Heland et al., 2001). In the first stripping coil HONO is quantitatively collected. Due to the acidic stripping solution, interfering species are collected less efficiently but in both channels. The true concentration of HONO is obtained by subtracting the inferences quantified in the second channel (in this study the average is 1 pptv, at most 5 pptv) from the total signal obtained from the first channel. For a more detailed description of LOPAP, see Heland et al. (2001). This correction of chemical interferences ascertained excellent agreement with the (absolute) differential optical absorption spectroscopy (DOAS) measurements, both in a smog chamber and under urban atmospheric conditions (Kleffmann et al., 2006). A possible interference from peroxynitric acid (HNO₄) has been proposed (Liao et al., 2006; Kerbrat et al., 2012; Legrand et al., 2014), but this will be insignificant at the high temperatures during the CYPHEX (CYprus PHotochemical EXperiment) campaign, at which HNO₄ is unstable. The stripping coils are temperature-controlled by a water-based thermostat, and the whole external sampling unit is shielded from sunlight by a small plastic housing. The reagents were all highpurity-grade chemicals, i.e., hydrochloric acid (37%, for analysis; Merck), sulfanilamide (for analysis, > 99 %; AppliChem) and N-(1-naphthyl)-ethylenediamine dihydrochloride (for analysis, >98%; AppliChem). For calibration Titrisol[®] 1000 mg NO₂⁻ (NaNO₂ in H₂O; Merck) was diluted to 0.0015 and $0.005 \text{ mg L}^{-1} \text{ NO}_2^{-1}$. For preparation all solutions were used, and for cleaning of the absorption tubes $18 M\Omega H_2O$ was used. The accuracy of the HONO measurements was 10%, based on the uncertainties of liquid and gas flow, concentration of calibration standard and regression of calibration.

NO and NO₂ measurements were made with a modified commercial chemiluminescence detector (CLD 790 SR), originally manufactured by ECO Physics (Duernten, Switzerland). The two-channel CLD based on the chemiluminescence of the reaction between NO and O₃ was used for measurements of NO and NO₂. NO₂ was measured as NO using a photolytic converter from Droplet Measurement Technologies (Boulder, USA). In the current study, data were obtained at a time resolution of 5 s. The CLD detection limits (determined by continuously measuring zero air at the measuring site) for NO and NO₂ measurements were 5 and 20 pptv, respectively for an integration period of 5 s. O₃ was measured with a standard UV photometric detector (Model 49, Thermo Environmental Instruments Inc.) with a detection limit of 1 ppb. Data are reported for an integration period of 60 s. The total uncertainties (2σ) for the measurements of NO, NO₂ and O₃ were determined to be 20, 30 and 5 %, respectively, based on the reproducibility of in-field background measurements, calibrations, the uncertainties of the standards and the conversion efficiency of the photolytic converter (Li et al., 2015).

OH and HO₂ radicals were measured using the HydrOxyl Radical measurement Unit based on fluorescence Spectroscopy (HORUS) setup developed at the Max Planck Institute for Chemistry (Mainz, Germany). HORUS is based on laser-induced fluorescence-fluorescence assay by gas expansion (LIF-FAGE) technique, wherein OH radicals are selectively excited at low pressure by pulsed UV light at around 308 nm, and the resulting fluorescence of OH is detected using gated microchannel plate (MCP) detectors (Martinez et al., 2010; Hens et al., 2014). The HORUS instrument had an inlet pre-injector (IPI) (Novelli et al., 2014) which allows the periodic addition of propane to scavenge the atmospheric OH radicals. This procedure allows the removal of potential interference species. HO2 is estimated by converting atmospheric HO₂ into OH using NO and detecting the additional OH formed. The instrument is calibrated by measuring signals from known amounts of OH and HO₂ generated by photolysis of water vapor in humidified zero air. The accuracy (2σ) of the OH measurements was 29%, and the precision (1σ) was 4.8×10^5 molecules cm⁻³.

Photolysis frequencies were determined using a spectroradiometer (Metcon GmbH) with a single monochromator and 512 pixel CCD array as a detector (275–640 nm). The thermostatted monochromator–detector unit was attached via a 10 m optical fiber to a 2- Π integrating hemispheric quartz dome. The spectroradiometer was calibrated prior to the campaign using a 1000 W National Institute of Standards and Technology (NIST) traceable irradiance standard. *J* values were calculated using molecular parameters recommended by the IUPAC and NASA evaluation panels (Sander et al., 2011; IUPAC, 2015). The *J* value for HONO was not corrected for upwelling UV radiation and is estimated to have an uncertainty of ~ 10 % (Bohn et al., 2008).

Aerosol measurements were also performed during the campaign. In this study particulate nitrate and aerosol surface data were used. These were detected by high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS, Aero-dyne Research Inc., Billerica, MA USA), and scanning mobility particle sizer (SMPS 3936, TSI, Shoreview, MN USA)



Figure 1. Map of location: the red star shows the location of Ineia and the measuring site. The four red points mark the main cities of Cyprus: Nicosia, Larnaca, Limassol and Paphos (clockwise ordering). Map produced by the Cartographic Research Lab University of Alabama; map of Cyprus: Google Maps.

and aerodynamic particle sizer (APS 3321, TSI), respectively. The mobility- and aerodynamics-based size distributions were combined based on the algorithm proposed by Khlystov et al. (2004).

The VOCs including α -pinene, β -pinene, isoprene, Δ 3carene, limonene and DMS (dimethyl sulfide) were detected by a commercial gas chromatography-mass spectrometry (GC-MS) system (MSD 5973; Agilent Technologies GmbH) coupled with an air sampler and a thermal desorber unit (Markes International GmbH). The VOCs were trapped at 30 °C on a low-dead-volume quartz cold trap (U-T15ATA; Markes International GmbH) filled with two-bed sorbent (Tenax TA and Carbograph I). The cold trap was heated to 320 °C, and the sample was transferred to a 30 m GC column (DB-624, 0.25 mm I.D., 1.4 µm film; J&W Scientific). The temperature of the GC oven was programmed to be stable at 40 °C for 5 min and then rise at a rate of $5 °C min^{-1}$ up to 140 °C. Thereafter, the rate was increased to 40 °C min⁻¹ up to 230 °C, where it was stabilized for 3 min. Each sample was taken every 45 min; calibrations, using a commercial gas standard mixture (National Physical Laboratory, UK), were performed every 8-12 samples.

Carbon monoxide was measured by infrared absorption spectroscopy using a room temperature quantum cascade laser at a time resolution of 1 s. Data are reported as 60 s averages with a total uncertainty of ~ 10 %, mainly determined by the uncertainty of the NIST standard used (Li et al., 2015).

Meteorological parameters (temperature, relative humidity (RH), wind speed and direction, pressure, solar radiation, precipitation) were detected by the weather station Vantage Pro2 from Davis Instruments.

Besides GC-MS all other operating instruments had time resolutions between 20 s and 5 min. For most analyses in this study the data were averaged to 10 min. When GC-MS data were included in the evaluation, 1 h averaged data were used.

3 Site description

Cyprus is a 9251 km² island in the southeast Mediterranean Sea (Fig. 1). The measuring site was located on a military compound in Ineia, Cyprus (34.9638° N, 32.3778° E), about 600 m above sea level and approximately 5.5–8 km from the coastline (main wind direction: W–SW). The field site is characterized by light vegetation cover, mainly comprising small shrubs like *Pistacia lentiscus, Sarcopoterium spinosum* and *Nerium oleander*; herbs like *Inula viscosa* and *Foeniculum vulgare*; and few typical Mediterranean trees like *Olea europaea, Pinus* sp. and *Ceratonia siliqua*. The area within a radius of about 15 km around the station is only weakly populated. Paphos (88 266 citizens) is located 20 km south of the field site; Limassol (235 000), Nicosia (325 756) and



Figure 2. Airflow conditions during the CYPHEX campaign: (a) measured local wind direction, (b) back trajectories calculated with NOAA Hysplit model showing examples for the two main air mass origins (48 h, UTC = LT - 3 h).

Larnaca (143 367) are 70, 90 and 110 km to the E-SE, respectively (population data according to statistical service of the Republic of Cyprus, http://www.cystat.gov.cy, census of population October 2011). During the campaign (7 July-3 August 2014), clear-sky conditions prevailed and occasionally clouds skimmed the site. No rain was observed, but the elevated field site was impacted by fog during nighttime and early morning due to adiabatic cooling of ascending marine humid air masses. Temperature ranged from 18 to 28 °C. Within the main local wind direction of SW (Fig. 2a) there was no direct anthropogenic influence, resulting in clean humid air from the sea. Analysis of 48 h back trajectories showed mainly two source regions of air mass origin (Fig. 2b). For approximately half (46%) of the campaign the air masses came from west of Cyprus, spending most of their time over the Mediterranean Sea prior to arriving at the site. During the remaining half of the campaign air masses originated from north of Cyprus, from eastern European countries (Turkey, Bulgaria, Rumania, Ukraine and Russia). Westerly air masses have been shown to exhibit lower concentration of gaseous and aerosol pollutants than the predominant northerly air masses that typically reach the site (Kleanthous et al., 2014). They spent more time over continental terrestrial surface and were likely to be additionally affected by biomass burning events detected in eastern Europe within the measurement periods (FIRMS, MODIS, web fire mapper, Fig. S1 in the Supplement). Previous back-trajectory studies in the eastern Mediterranean support this assumption (Kleanthous et al., 2014; Pikridas et al., 2010).

Most of the time the advected air mass was loaded with high humidity as a result of sea breeze circulation. Two periods of about 4 days with lower relative humidity occurred. These two situations will be contrasted below.

4 Results

The concentrations of HONO and other atmospheric trace gases as well as meteorological conditions observed on Cyprus from 7 July to 3 August 2014 are shown in Fig. 3. In general, low trace gas mixing ratios were indicative of clean marine atmospheric boundary conditions, as pollutants are oxidized by OH during the relatively long air transport time over the Mediterranean Sea (more than 30 h), and without significant impact of direct anthropogenic emissions.



Figure 3. Measured variables during the whole campaign from 7 July to 4 August 2014. (a) Meteorological data (temperature, T; relative humidity, RH; wind direction, wd; wind speed, ws), O₃ and CO indicate stable conditions; in the lower panel the bar indicates the air mass origin: bright blue represents westerly, while the brownish color represents northerly. (b) Observed mixing ratios of HONO, NO₂ and NO, and the photolysis frequency J_{HONO} and the HONO / NO_x ratio. The yellow and blue boxes reflect the dry and the humid periods, respectively.

Ambient HONO mixing ratios ranged from below detection limit (<4 pptv) to above 300 pptv. Daily average HONO was 35 pptv (± 25 pptv; 1σ standard deviation). The daily average NO₂ and NO mixing ratios were 140 ± 115 and 20 ± 35 pptv, respectively, but showed intermittent peaks up to 50 ppbv when sampling air was streamed from the diesel generator used to power the station, from the access route or the parking lot by local winds (easterly, Fig. S2). These incidents, which account for 4% of the campaign time, were classified as local air pollution events and were omitted from analysis. Mean O3 and CO mixing ratios were 72 ± 12 ppb and 98 ± 11 ppbv, respectively. OH radicals ranged from below detection limit $(1 \times 10^5 \text{ molecules cm}^{-3})$ during nighttime to 8×10^6 molecules cm⁻³ during daytime (see Fig. S3). Daytime HO₂ / OH ratio ranged from 100 to 150. The mixing ratios of NO₂, O₃ and CO varied in unison and were significantly (p < 0.05) higher during periods when air masses originated from eastern Europe (brownish bar in Fig. 3a, lower panel), indicative of air pollution and shorter transport times compared to western Europe (NO₂: northerly: 144 ± 130 pptv, westerly: 127 ± 106 pptv; O₃: northerly: 74 ± 11 ppbv, westerly: 66 ± 12 ppbv; CO: northerly: 101 ± 9 ppbv, westerly: 90 ± 10 ppbv). In contrast, NO and HONO mixing ratios were slightly higher when air masses came from western Europe and over the sea (NO: northerly: 17 ± 35 pptv, westerly: 20 ± 44 pptv; HONO: northerly: 32 ± 26 pptv, westerly: 38 ± 22 pptv).

Besides two different air mass origins, two periods with different behavior of relative humidity were identified, as illustrated by blue and yellow boxes in Fig. 3a and b. In both periods we found northerly and westerly air mass origins. The diel profiles of trace gas mixing ratios and meteorological variables of the humid period (blue box) are shown in Fig. 4a, and the ones of the dry period (yellow box) in Fig. 4b. During the drier period HONO concentrations were stable and low (6 pptv) during nighttime, while mean nighttime HONO mixing ratios during the humid period (Fig. 4a) showed an expected slow increase of about 20 pptv (from 20 to 40 pptv), as anticipated from heterogeneous production and accumulation within a nocturnal boundary layer characterized by a stable stratification and low wind speed (Acker et al., 2005; Su et al., 2008b; Li et al., 2012). During both periods, but more pronounced in the drier period, HONO rapidly increased by a factor of 2 within 2h after sunrise and then slowly decreased until sunset. Similar profiles were also observed for other trace gases, like isoprene or DMS, which are transported in upslope winds. Strong HONO morning peaks and high daytime mixing ratios suggest a strong daytime source, compensating the short atmospheric lifetime (15 min) caused by fast photolysis.

Mean NO mixing ratios were close to the detection limit (5 pptv) at night and increased after sunrise (06:00 local time, LT) to mean values of 60 pptv (peak 150 pptv) at 09:00 LT, prior to declining for the rest of the day until sunset (20:00 LT). In the absence of local NO sources low night-time values are a result of the conversion of NO to NO₂ by O₃, which was continuously high (Hosaynali Beygi et al., 2011). The diel profiles of NO mixing ratios followed closely those of HONO mixing ratios. This similarity and their dependency on relative humidity are suggestive of a common source for both reactive nitrogen species.

 NO_2 mixing ratios were somewhat lower during nighttime, but in general the diel variability remained in a narrow range between 100 and 200 pptv. Likewise, the diel courses of O_3 and CO mixing ratios revealed relatively low day–night variability in a range of 65–75 and 90–100 ppb, respectively.



Figure 4. Diel variation of meteorological data (temperature, *T*; relative humidity, RH), NO and NO₂ mixing ratios, the photolysis rate for HONO J_{HONO} and HONO mixing ratios (pink: measured; violet: daytime photostationary state (PSS); grey: nighttime heterogeneous NO₂ conversion) and HONO / NO_x ratio for (**a**) average for period when RH was above 60 % (blue box in Fig. 3) and (**b**) average for dry period when RH was below 60 % (yellow box in Fig. 3). Error bars represent standard deviation of diel mean.

5 Discussion

Low-NO_x conditions at this remote field site in photochemically aged marine air were found to be an ideal prerequisite to trace as yet undefined local HONO sources. On Cyprus, diel profiles of HONO showed peak values in the late morning and persistently high mixing ratios during daytime, as has been reported for some other remote regions (Acker et al., 2006a; Zhou et al., 2007; Huang et al., 2002). This is not the case for rural and urban sites, where atmospheric HONO mixing ratios are normally observed to continuously build up during nighttime, presumably due to heterogeneous reactions involving NO_x and decline in the morning due to strong photodissociation (e.g., Elshorbany et al., 2012, and references therein).

The diel HONO / NO_x ratio (Fig. 4a, b, third panel) shows consistently high values during the humid period (Fig. 4a) and significant diel variation for the dry case (Fig. 4b) with higher values during daytime. The ratio (average of 0.33 and peak values greater than 2) is higher than that reported for most other regions, suggesting a strong impact of local HONO sources. Elshorbany et al. (2012) investigated data from 15 different urban and rural field measurement campaigns around the globe, and came up with a robust representative mean atmospheric HONO / NO_x ratio as low as 0.02. However, high values were observed at remote mountain sites, with mean values of 0.23 (up to ≈ 0.5 in the late morning; Zhou et al., 2007) or 0.2–0.4 at remote Arctic/polar sites (Li, 1994; Zhou et al., 2001; Beine et al., 2001; Jacobi et al., 2004; Amoroso et al., 2010). Legrand et al. (2014) observed HONO / NO_x ratios between 0.27 and 0.93 during experiments with irradiated Antarctic snow, depending on radiation wavelength, temperature and nitrate content. Elevated HONO / NO_x ratios at low NO_x levels show the importance of HONO formation mechanisms other than heterogeneous NO_x reactions.

5.1 Nighttime HONO accumulation

Between 18:30 and 07:30 LT HONO has an atmospheric lifetime of more than 45 min and [OH] is low, just about 1×10^5 molecules cm⁻³, so that the calculation of HONO at photostationary state [HONO]_{pss} (Reactions R1–R3) at night is not appropriate. Instead, nighttime HONO concentrations can be estimated due to heterogeneous reaction of NO₂ described in Eq. (1) (Alicke et al., 2002, 2003; Su et al., 2008b; Sörgel et al., 2011b). Three studies in different environments from a rural forest region in eastern Germany (Sörgel et al., 2011b) and a non-urban site in the Pearl River Delta, China (Su et al., 2008b), to an urban, polluted site in Beijing (Spataro et al., 2013) found a conversion rate of about 1.6 % h⁻¹ (1.1–1.8 % h⁻¹).

 $[\text{HONO}]_{\text{het}} = [\text{HONO}]_{\text{evening}} + 0.016 \,\text{h}^{-1} [\text{NO}_2] \Delta t \tag{1}$

 $[HONO]_{het}$ denotes the accumulation of HONO by heterogeneous conversion of NO₂, $[HONO]_{evening}$ the measured HONO concentration at 20:30 LT, [NO₂] the measured average NO₂ concentration between 20:30 and 07:30 LT, and Δt time span in hours.

Measured and calculated HONO mixing ratios are compared in Fig. 4 (upper panel). During the humid period, during nighttime the estimated (according Eq. 1; Fig. 4a, upper panel, grey line) and observed HONO mixing ratios are in good agreement ($R^2 = 0.9$). During the drier period the observed HONO mixing ratios were lower than the ones calculated with a NO₂ conversion rate of 1.6 % h⁻¹. Here the approach for the nighttime conversion frequency by, e.g., Alicke et al. (2002, 2003), Su et al. (2008b) or Sörgel et al. (2011b) (rate = $\frac{\text{HONO}_{t2} - \text{HONO}_{t1}}{\Delta t \cdot \text{NO}_2}$) was used. The 7-day average conversion rate for the dry nights was 0.36 % h⁻¹ (Fig. 4b, upper panel, black line), comparable to results of Kleffmann et al. (2003) reporting a conversion rate of $6 \times 10^{-7} \text{ s}^{-1}$ (0.22 % h⁻¹) for rural forested land in Germany.

As already mentioned above, it is apparent that HONO mixing ratios under low-RH conditions during nighttime were much lower than under humid conditions, and HONO morning peaks were most pronounced (compare Fig. 4a and b: humid/dry). HONO (Donaldson et al., 2014a) and NO₂ (Wang et al., 2012; Liu et al., 2015) uptake coefficients have recently been reported to be much stronger for dry soil and at low RH, respectively, which is in line with HONO on Cyprus being close to the detection limit on nights with low relative humidity. On the other hand, it has been shown on glass and on soil proxies that the yield of HONO formation from NO₂ on surfaces is low under dry conditions but sharply increases at RH > 30 % (Liu et al., 2015) or > 60 % (Finlayson-Pitts et al., 2003). On Cyprus the strong morning HONO peaks after dry nights were accompanied by an increase in relative humidity from 40 to 80%. Deposited and accumulated NO₂ on dry soil surfaces could be released as HONO at high rates under elevated-RH conditions. In contrast, in a humid regime HONO mixing ratios were continuously high during nighttime and showed less pronounced morning peaks, suggesting lower nighttime deposition of NO₂ and lower HONO emissions in the morning, respectively.

As morning HONO peak mixing ratios were most pronounced after dry nights on Cyprus, our observations are to some extent contradictory to earlier results that have proposed that dew formation on the ground surface may be responsible for HONO nighttime accumulation in the aqueous phase, followed by release from this reservoir after dew evaporation the next morning (Zhou et al., 2002a; Rubio et al., 2002; He et al., 2006). We cannot rule out that the latter could have contributed to nighttime accumulation of HONO during humid conditions, as we had no means to measure dew formation at the site, and high daytime HONO mixing ratios were observed under all humidity regimes. However, kinetic models of competitive adsorption of trace gases and water onto particle surfaces predict exchange behavior explicitly distinct from the liquid phase (Donaldson et al., 2014a). The nitrogen composition in thin water films (few water molecular monolayers) is complex, including HONO, NO, HNO₃, water–nitric acid complexes, NO_2^+ and N_2O_4 (Finlayson-Pitts et al., 2003). With only small amounts of surface-bound water, nitric acid is largely undissociated HNO₃ and is assumed to be stabilized upon formation of the HNO₃–H₂O complexes (hydrates), which have unique reactivity compared to nitric acid water aqueous solutions, where it is dissociated H⁺ and NO_3^- ions (Finlayson-Pitts et al., 2003). Likewise, HONO formation rates in surface-bound water are about 4 orders of magnitude larger than expected for the aqueous-phase reaction (Pitts et al., 1984).

Diel HONO profiles very similar to those on Cyprus with a late-morning maximum and late-afternoon/early-evening minimum have been observed at the Meteorological Observatory Hohenpeissenberg, a mountain-top site in Germany (Acker et al., 2006a) and by Zhou et al. (2007) at the summit of Whiteface Mountain in New York State. For the latter study, formation of dew could be ruled out as relative humidity was mostly well below saturation. Zhou et al. (2007) argued that the high HONO mixing ratios during morning and late morning can be explained by mountain up-slope flow of polluted air from the cities at the foot of the mountain that results from ground surface heating. On Cyprus the sea breeze, driven by the growing difference between sea and soil surface temperature, brings air to the site which interacted with the soil surface and vegetation and is loaded by respective trace gas emissions. This is endorsed by the simultaneous increase of DMS and isoprene, markers for transportation of marine air and emission by vegetation. In the late afternoon, when the surface cools, down-welling air from aloft would dominate, being less influenced by ground surface processes. Zhou et al. (2007) could show that noontime HONO mixing ratios and average NO_v during the previous 24 h period were strongly correlated, much better than instantaneous HONO / NO_y or HONO / NO_x, which is in line with N accumulation on soil surfaces as discussed above.

5.2 Daytime HONO budget

During daytime (07:30 to 18:00 LT, with HONO lifetime being between 10 and 30 min), [HONO]_{PSS}, the photostationary HONO concentration resulting from gas-phase chemistry, can be calculated according to Eq. (2) (Kleffmann et al., 2005):

$$[\text{HONO}]_{\text{PSS}} = \frac{k_1 [\text{OH}] [\text{NO}]}{k_2 [\text{OH}] + J_{\text{HONO}}},$$
(2)

where k_1 and k_2 are the temperature-dependent rate constants for the gas-phase HONO formation from NO and OH and the loss of HONO by reaction of HONO and OH, respectively (Atkinson et al., 2004; e.g., at 23.0 °C a typical temperature during this study $k_1 \approx 1.36 \times 10^{-11} \text{ cm}^3 \text{ s}^{-1}$; $k_2 \approx$ $6.01 \times 10^{-12} \text{ cm}^3 \text{ s}^{-1}$). J_{HONO} is the photolysis frequency of



Figure 5. HONO budget analysis for (a) the humid and (b) the dry period. S_{OH+NO} (black) stands for the formation rate of HONO via the reaction of NO and OH, $S_{Het_NO_2}$ (yellow) is the formation rate for the heterogeneous reaction of NO₂ (conversion rate $a = 1.6 \% h^{-1}$; $b = 0.36 \% h^{-1}$), L_{phot} (green) and $L_{OH+HONO}$ (blue) are the loss rates via photolysis, and the reaction with OH and $S_{unknown}$ is the unknown source. Error bars indicate standard deviation of diel mean.

HONO, which was measured with a spectroradiometer. [NO] is the observed NO concentration. Since OH data were available only on a few days, diel variations of [OH] were averaged (see Fig. S3).

As has been previously established by many other studies (Su et al., 2008a; Michoud et al., 2014; Sörgel et al., 2011a), homogeneous gas-phase chemistry alone fails to reflect observed HONO mixing ratios. Observed daytime values were up to 30 times higher than calculated based on PSS, indicating strong additional local daytime sources of HONO. Lee et al. (2013) argue that the HONO PSS assumption might overestimate the strength of any unidentified source if the transport time from nearby NO_x emission sources to the measurement site is less than the time required for HONO to reach PSS. In this study, the missing source was calculated according to Su et al. (2008a) (Eq. 3), where PSS was not assumed. Also in our measurements, dHONO/dt was not equal to 0, as HONO was not at PSS.

$$S_{\text{HONO}} = J_{\text{HONO}}[\text{HONO}] + k_2[\text{OH}][\text{HONO}]$$
$$-k_1[\text{OH}][\text{NO}] - k_{\text{het}}[\text{NO}_2] + \frac{\Delta[\text{HONO}]}{\Delta t}$$
(3)

[HONO] is the measured HONO concentration and k_{het} the heterogeneous conversion rate of NO₂ to HONO, which was discussed above to be 1.6 % h⁻¹ during the wet period and 0.36 % h⁻¹ during the dry period. Δ [HONO] / Δt is the observed change of HONO concentration unequal to 0. The uncertainty of the calculated missing source S_{HONO} was estimated to be about 16 % based on the Gaussian error propagation of instrument uncertainties of HONO, NO, NO₂, J and OH.

Nevertheless, at the study site of Cyprus, the mean upwind distance between the measurement site and the coastline was about 6 km, and the mean wind velocity was about 3 m s^{-1} . Accordingly, the respective air mass travel time over land is estimated to be about half an hour, which is somewhat

longer than the daytime lifetime of HONO and might provide enough time for the equilibrium processes. Furthermore and in strong contrast to Lee et al. (2013), at the Cyprus site the concentrations of HONO precursors (NO and OH) were extremely low, far too low to explain the observed HONO concentrations. In the late morning (around 10:00 LT) the unknown source was at its maximum, with peak production rates of up to 3.4×10^6 molecules cm⁻³ s¹ and a daytime average of about 1.3×10^6 cm⁻³ s⁻¹, which is in good agreement with other studies at rural sites, like a mountain site at Hohenpeissenberg ($(3 \pm 1) \times 10^6$ cm⁻³ s⁻¹, at NO_x \approx 2 ppbv; Acker et al., 2006a), a deciduous forest site in Jülich $(3.45 \times 10^6 \text{ molecules cm}^{-3} \text{ s}^{-1}, \text{ at NO} \approx 250 \text{ pptv}; \text{ Kleff-}$ mann et al., 2005) and a pine forest site in southwest Spain 0.74×10^6 molecules cm⁻³ s⁻¹, at NO_x ≈ 1.5 ppby; Soergel et al., 2011a) but smaller than at urban sites in Houston $(4-6 \times 10^6 \text{ cm}^{-3} \text{ s}^{-1})$, at NO_x \approx 6 ppby; Wong et al., 2012), Beijing $(7 \times 10^6 \text{ cm}^{-3} \text{ s}^{-1})$, at NO_x ≈ 15 ppby; Yang et al., 2014) and southern China $(5.25 \pm 3.75 \times 10^6 \text{ cm}^{-3} \text{ s}^{-1})$, at NO_x \approx 20 ppbv (Li et al., 2012), or 1–4 \times 10⁷ cm⁻³ s⁻¹, at $NO_x \approx 35$ ppbv (Su et al., 2008a)).

The contributions of gas-phase reactions and the heterogeneous reaction of NO₂ (conversion rate *a* 1.6 % h⁻¹ and *b* 0.36 % h⁻¹) to the HONO budget are illustrated in Fig. 5 exemplarily. For both periods the contributions are quiet similar; just the absolute values are different. To compensate the strong loss via photolysis, a comparably strong unknown source is necessary as the heterogeneous NO₂ conversion or the gas-phase reaction of OH and NO is insignificant.

In polluted regions with moderate to high NO_x concentrations, HONO sources have often been linked with [NO₂] or [NO_x] (Acker et al., 2005; Li et al., 2012; Levy et al., 2014; Sörgel et al., 2011a; Wentzel et al., 2010). Under the prevailing low-NO_x conditions during CYPHEX (< 250 pptv), correlation analysis (see Table 1) of S_{HONO} with [NO₂] ($R^2 =$ 0.50) and [NO₂] · RH ($R^2 = 0.51$) indicate no significant im-

		During the whole campaign							
			Time-of-day average						
			HONO	S _{HONO}	HONO	S_{HONO}			
	Г		0.006	0.125	0.488	0.214	-		
]	RH		0.077	0.005 ^d	0.092	0.103			
]	Heat flux		0.261	0.300	0.617 ^c	0.585 ^c			
	$J_{\rm NO_2}$		0.263	0.395	0.718 ^b	0.672 ^b			
1	NO		0.242	0.154	0.857 ^a	0.600 ^c			
I	NO ₂		0.052	0.078	0.620^{c}	0.496			
I	$NO_2 \cdot RH$		0.126	0.111	0.638 ^c	0.505 ^c			
]	$NO_2 \cdot RH \cdot aerosol$	0.095	0.092	0.256	0.579 ^c				
]	$NO_2 \cdot J$	0.191	0.164	0.828 ^a	0.813 ^a				
]	$NO_2 \cdot RH \cdot J$		0.266	0.221	0.850 ^a	0.807 ^a			
]	$NO_2 \cdot RH \cdot J \cdot aerosol surface$		0.221	0.204	0.806 ^a	0.814 ^a			
	S _{NO}			0.012		-0.015^{d}			
	During the			umid period Durin			the dry period		
	-		Time-of-d	ay average		Time-of-day average			
	HONO	S _{HONO}	HONO	S_{HONO}	HONO	S _{HONO}	HONO	S _{HONO}	
Т	0.006	0.116	0.031	0.123	0.120	0.016	0.453	-0.004	
RH	0.000	0.081 ^d	0.010 ^d	0.146 ^d	0.374	0.193	0.730 ^b	0.603 ^c	
Heat flux	0.110	0.243	0.184	0.591 ^c	0.502 ^c	0.335	0.685 ^b	0.634 ^c	
$J_{\rm NO_2}$	0.150	0.465	0.245	0.669 ^b	0.678 ^b	0.320	0.829 ^a	0.664 ^b	
NO	0.168	0.135	0.418	0.650 ^b	0.487	0.301	0.730 ^b	0.409	
NO ₂	0.066	0.065	0.300	0.267	0.037	0.003 ^d	0.619 ^c	0.174	
$NO_2 \cdot RH$	0.084	0.048	0.294	0.171	0.161	0.010	0.714 ^b	0.456	
$NO_2 \cdot RH \cdot aerosol surface$ 0.047		0.072	0.111	0.250	0.241	0.085	0.557 ^c	0.551 ^c	
$NO_2 \cdot J$	0.214	0.261	0.427	0.845 ^a	0.358	0.016	0.872 ^a	0.603 ^c	
$NO_2 \cdot RH \cdot J$	0.231	0.244	0.467	0.775 ^b	0.434	0.068	0.820 ^a	0.703 ^b	
$NO_2 \cdot RH \cdot J \cdot aerosol surface 0.140$ (0.152	0.465	0.795 ^b	0.414	0.130	0.664 ^b	0.631 ^c	
S _{NO}		0.294		0.720 ^b		0.059		0.094	

Table 1. Linear correlation factors (Pearson correlation, R^2) of HONO and the unknown source S_{HONO} to meteorological factors and different NO_x parameters.

^a Highly correlated: $R^2 > 0.8$ (in bold font). ^b Moderately correlated: $0.65 < R^2 < 0.8$ (in italic font). ^c Poorly correlated: $0.5 < R^2 < 0.65$ (in normal font).

^d Anti-correlated.

pact of instantaneous heterogeneous formation of HONO from NO₂. Better correlations of S_{HONO} with J_{NO_2} ($R^2 =$ 0.67) and $J_{NO_2} \cdot [NO_2]$ ($R^2 = 0.82$) indicate a photo-induced conversion of NO2 to HONO as already suggested by George et al. (2005) or Stemmler et al. (2006, 2007). Lee et al. (2016) found even lower correlation with [NO₂] ($R^2 = 0.0001$) but similar good correlation with $J_{NO_2} \cdot [NO_2]$ ($R^2 = 0.70$) at an urban background site in London. Other light-dependent reactions such as the photolysis of nitrate might additionally contribute to high daytime HONO. It is unlikely that aerosol surfaces played an important role in heterogeneous conversion of NO2 as the mean observed aerosol surface concentration was only about $300 \,\mu\text{m}^2 \,\text{cm}^{-3}$. Based on a formula for photo-enhanced conversion of NO2 on humic acid aerosols which was derived by Stemmler et al. (2007), a HONO formation rate of only 5.1×10^2 molecules cm⁻³ s⁻¹ can be estimated. Likewise, Sörgel et al. (2015) showed that HONO fluxes from light-activated reactions of NO2 on humic acid surfaces at low NO₂ levels (< 1 ppb and thus comparable to concentrations observed in this study) saturated at around 0.0125 nmol m⁻² s⁻¹. Therefore heterogeneous aerosol surface reactions can be neglected as HONO sources at the prevailing low NO_x levels.

Likewise, the nitrate concentrations of highly acidic marine aerosols particulate matter as measured by HR-ToF-AMS (PM1 fraction, mean of 0.075 µg m⁻³) were too low to account for significant photolytic HONO production $(1.7 \times 10^2 \text{ molecules cm}^{-3} \text{ s}^{-1} \text{ or } 0.01 \% \text{ of } S_{\text{HONO}})$ calculated by Eq. (4):

$$S_{\text{photo}_\text{NO}_3^-} = [\overline{\text{NO}_3^-}] \cdot J_{\text{NO}_3^-}, \tag{4}$$

with $S_{\text{photo}_NO_3^-}$ being the source strength of HONO by photolysis of nitrate, $[\overline{NO_3^-}]$ the mean particulate nitrate concentration and $J_{NO_3^-}$ the photolysis frequency of nitrate (aqueous) at noon (3 × 10⁻⁷ s⁻¹; Jankowski et al., 1999).

Recently an enhancement of the photolysis frequency of particulate nitrate relative to gaseous or aqueous nitrate was found (Ye et al., 2016). But even with this enhanced rate of $2 \times 10^{-4} \text{ s}^{-1}$ no more than 1.1×10^5 molecules cm⁻³ s⁻¹ (8 % of *S*_{HONO}) HONO would be produced.

5.3 Common daytime source of HONO and NO

During CYPHEX, good correlation was found between [HONO] or S_{HONO} and [NO] ($R^2 = 0.86$ and 0.60, respectively), indicating that both may have a common source. A missing source of NO can be calculated as shown in Eq. (5).

$$S_{\rm NO} = k_1 [\rm OH][\rm NO] + k_3 [\rm HO_2][\rm NO] + k_4 [\rm O_3][\rm NO] + k_5 [\rm RO_2][\rm NO] - J_{\rm NO_2}[\rm NO_2] - J_{\rm HONO}[\rm HONO] + \frac{\Delta[\rm NO]}{\Delta t}$$
(5)

 k_3 and k_4 are the temperature-dependent rate constants for the reaction of NO with HO₂ and O₃, respectively (Atkinson et al., 2004; at 23 °C: $k_3 \approx 8.96 \times 10^{-12}$ cm³ s⁻¹; $k_4 \approx 1.68 \times$ 10^{-14} cm³ s⁻¹); k_5 is the rate constant for the reaction of NO and organic peroxy radicals which was assumed to be the same as for the reaction NO + CH₃O₂ (7.7 × 10⁻¹² cm³ s⁻¹ at 298 K; Ren et al., 2010; Sander et al., 2011). Like [OH], [HO₂] was also measured only on a few days, and therefore mean diel data were used (Fig. S3). Total [RO₂] was estimated to be maximum $1.6 \cdot$ [HO₂] (Ren et al., 2010; Hens et al., 2014). Using a RO₂ / HO₂ ratio of 1.2, the absolute values of *S*_{NO} are reduced by 0.3 to 5.5 %. The budget analysis for NO for both humidity regimes is illustrated in Fig. S4.

For NO_x , an unexpected deviation from the PSS, or Leighton ratio, of clean marine boundary layer air has been observed previously, invoking a hitherto unknown NO sink, or pathway for NO to NO₂ oxidation, other than reactions with OH, HO₂, O₃ and organic peroxides (Hosaynali Beygi et al., 2011). On Cyprus, two different atmospheric humidity regimes can be differentiated. Under dry conditions (RH < 70 %, yellow boxes in Fig. 3) and higher NO_x concentrations (> 150 pptv) S_{NO} is negative, implying a net NO sink of up to 6.4×10^7 molecules cm⁻³ s⁻¹ resembling the abovementioned PSS deviations in remote marine air masses (see Figs. 6 and 7). However, during humid conditions (RH > 70,blue boxes in Fig. 3) $S_{\rm NO}$ was positive with values of up to 5.1×10^7 molecules cm⁻³ s⁻¹. Due to low and invariant acetonitrile levels, anthropogenic activity and local biomass burning can be excluded as an NO source at this specific site. A net NO source during humid conditions is assumed to result from (biogenic) NO emission from soil. As shown in Fig. 8, the S_{HONO} and S_{NO} (time-of-day average, excluding 3 days as there are transition days (25 July and 2 August) or the RH changed too quickly (15 July)) were highly correlated ($R^2 = 0.72$), indicative of both reactive N compounds being emitted from the same local source. Both HONO and NO have been reported to be released from soil, with a strong dependency on soil water content (Su et al., 2011; Oswald et



Figure 6. NO₂ (color-coded) and RH dependence of the sources of NO (S_{NO}) and HONO (S_{HONO}).

al., 2013; Mamtimin et al., 2016). The (dry-state) soil humidification threshold level for NO emission is reported to be somewhat higher than for HONO (Oswald et al., 2013), which might explain why a net NO source was preferentially calculated for higher-relative-humidity conditions, while for HONO a daytime source under all humidity regimes prevailing during the campaign was found. Mamtimin et al. (2016) investigated HONO and NO emissions of natural desert soil and with grapes or cotton cultivation soils in an oasis in the Taklamakan Desert in the Xinjiang region in China. After irrigation they did not find direct emission, but when the soil had almost dried out (gravimetric soil water content: 0.01-0.3) emissions up to $115 \text{ ng N m}^{-2} \text{ s}^{-1}$ were detected. In addition they observed soil-temperature-dependent emission of reactive nitrogen. Analyzing microbial surface communities from drylands, Weber et al. (2015) observed highly correlated NO-N and HONO-N emissions with Spearman rank correlation coefficients ranging between 0.75 and 0.99. In this study, NO and HONO emissions were observed in drying soils with water contents of 20-30 % water holding capacity.

Even though we cannot make firm conclusions regarding the exact mechanism of HONO formation, the abovementioned correlation analysis (and Table 1) reveal that the instantaneous heterogeneous NO2 conversion is not a significant HONO source. We propose that HONO is emitted from nitrogen compounds being accumulated on mountain slope soil surfaces produced either biologically by soil microbiota or from previously deposited NO_{ν} . This forms the major daytime HONO source responsible for morning concentration peaks and consistently high daytime mixing ratios at the Cyprus field site. While biological formation is assumed to be more relevant for humid conditions, physical NO_v accumulation can be assumed to be stronger under dry conditions, as uptake coefficients for a variety of trace gases were shown to be significantly higher for dry surfaces, among them NO₂ (Wang et al., 2012; Liu et al., 2015), HONO (Donaldson et



Figure 7. Diel profile of both unknown sources, S_{HONO} (**a**) and S_{NO} (**b**), for all data, humid (excluding transition days, 25 July and 2 August, and 15 July as RH conditions changed too quickly) and dry periods. Error bars indicate standard deviation of diel average.

al., 2014a) and HCHO (Li et al., 2016). The strongest HONO morning peaks observed after dry nights were accompanied by an increase in relative humidity driven by the sea breeze (Fig. 4b), so we consider HONO as being released preferentially under favorable humid conditions.

5.4 Primary OH production

Many studies showed high contribution of HONO photolysis to the OH budget (up to 30 % on average daily; Alicke et al., 2002; Ren et al., 2006). Here, the primary OH production rates are calculated based on the main OH-forming reactions, which are the photolysis of O_3 and subsequent reaction with water (Reactions R6, R7), the photolysis of HONO (Reaction R2) and the reaction of alkenes with ozone (Reaction R8).

$$O_3 \xrightarrow{hv(< 340 \text{ nm})} O\left(^1 \text{D}\right) + O_2 \tag{R6}$$

$$O(^{1}D) + H_{2}O \rightarrow 2OH$$
 (R7)

alkene +
$$O_3 \rightarrow OH$$
 + other products (R8)

Reaction rates were taken from Atkinson et al. (2004) and Atkinson (1997). The water pressure over water was calculated according to Murphy and Koop (2005). Reactions of $O(^{1}D)$ and HO₂ not forming OH are also considered. OH formation yields of the reactions of alkenes with O₃ were taken from Paulson et al. (1999). Photolysis rates (*J* values) and concentrations of relevant compounds were as measured on Cyprus. Isoprene, α -pinene, β -pinene, Δ 3-carene and limonene (VOC) were taken into account as the most relevant alkenes.

The results of this study are shown in Fig. 9. All three production routes show a clear diel profile with higher production rates during daytime. In the night only the reaction of alkenes with O₃ produced significant amounts of OH $(2 \times 10^4 \text{ molecules cm}^{-3} \text{ s}^{-1})$. With sunrise the other sources become more relevant. During daytime the photolysis of HONO generates about $1.5 \times$ 10^6 molecules OH cm⁻³ s⁻¹, which is about 10 times higher than the ozonolysis of alkenes at that time. The maximum OH production rate by O_3 photolysis during daytime is about 1.3×10^7 molecules cm⁻³ s⁻¹. In the morning (06:00– 08:00 LT) and evening hours (19:00-20:00 LT) the contribution of HONO photolysis to the primary OH production is on average 37 % (see Fig. 9b) with peak values of 65 %, which is much higher than the contribution of O_3 photolysis at that time. During the rest of the day the contribution of HONO decreases to 12 %. At noon the most dominant OH source is the photolysis of O_3 (more than 80 %), while the contribution of the ozonolysis of alkenes is almost negligible (1-2%). A complete and detailed HO_x budget analysis with CYPHEX data will be published soon.

6 Conclusion

Nitrous acid was found in low concentrations on the east Mediterranean island of Cyprus during summer 2014. Daytime concentrations were much higher than during the night and about 30 times higher than would be expected by budget analysis based on photostationary state. The unknown source was calculated to be about 1.9×10^6 molecules cm⁻³ s⁻¹ around noon. Low NO_x concentrations, high HONO / NO_x



Figure 8. Correlation of S_{HONO} to light-induced NO₂ reaction (for both periods; humid: blue triangle; dry: orange square), to NO and S_{NO} (only for humid period, excluding the transition days – 25 July and 2 August – and the day with quickly changing RH – 15 July); time-of-day-average data were used.



Figure 9. Average diel pattern of primary OH production from HONO, O_3 and VOC, shown as (a) production rate and (b) percentage contributions to primary OH production.

ratio and low correlation between HONO and NO_2 indicate a local source which is independent of NO_2 . Heterogeneous reactions of NO_2 on aerosols play an insignificant role during daytime. Emission from soil, caused either by photolysis of nitrate or gas–soil partitioning of accumulated nitrite/nitrous acid, is supposed to have a higher impact on the HONO concentration during this campaign. Also the NO budget analysis showed a missing source in the humid period, which correlates well with the unknown source of HONO, indicating a common source. The most likely source of HONO and NO is the emission from soil.

Even though the HONO concentration is only in the lower pptv level, it has a high contribution to the primary OH production in the early morning and evening hours.

7 Data availability

Readers who are interested in the data should contact the authors: Hang Su (h.su@mpic.de) or Hannah Meusel (hannah.meusel@mpic.de).

The Supplement related to this article is available online at doi:10.5194/acp-16-14475-2016-supplement.

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